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L11 ANSWER 1 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
                          2007:1431771 CAPLUS <<LOGINID::20080328>>
ACCESSION NUMBER:
DOCUMENT NUMBER:
                          148:105767
                          Hexakis (3,6-anhydro)-tetrakis [2I,II,IV,V-0-(2-
                          ethoxyethyl)] derivatives of (3,6-anhydro)-a-
                          cyclodextrin exhibits novel cation affinities
                          and tensioactive properties on membranes
AUTHOR(S):
                          Debouzy, J. C.; Crouzier, D.; Gadelle, A.
CORPORATE SOURCE:
                          Biophysics Laboratory, Centre de recherches du service
                          de sante des armees, La Tronche, Fr.
                          Pharmazie (2007), 62(12), 892-899
                          CODEN: PHARAT; ISSN: 0031-7144
                          Govi-Verlag Pharmazeutischer Verlag GmbH
DOCUMENT TYPE:
LANGUAGE:
     The synthesis of hexakis (3,6-anhydro)-tetrakis[2I,II,IV,V-0-(2-
     ethoxyethyl)] cyclomaltohexaose (AEOE) was designed to obtain cation
     complexing properties. 1H NMR study showed ionic radius dependence of
     AEOE cation affinity, markedly observed for Cs+ and Rb+. Besides, AEOE was
     found hemolytic (HC50 = 9mM) and superficial tension measurements revealed
     pos. tensioactive properties. A 31P and 2HNMR study of phospholipid
     dispersions (dimyristoyl phosphatidyl choline, DMPC) in the presence of
     AEOE was performed; it was found that, beside the typical lineshape of
     phospholipid bilayers, two new NMR lines were detected in the presence of
     AEOE: (a) an isotropic line consistent with a detergent effect (b) another
     isotropic resonance of 1 Hz linewidth over phase transition temperature (298 K),
     indicating a true solubilization. Coupling constant measurements confirmed
     that the main conformation at the polar head group level was close to that
     observed in chloroform/methanol solution It was finally concluded that AEOE
     could form true solns. of DMPC, similarly to those induced by di-Et ether
     interactions with membranes, while giving soluble complexes.
                                THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                          38
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 2 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          2007:1220778 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                          Physicochemical properties and membrane interactions
                          of per(6-desoxy-6-halogenated) cyclodextri
AUTHOR(S):
                          Debouzy, J.-C.; Crouzier, D.; Gadelle, A.
CORPORATE SOURCE:
                          Unite de Biophysique, Centre de Recherches du Service
                          de Sante des Armees, La Tronche, F 38702, Fr.
SOURCE:
                          Annales Pharmaceutiques Françaises (2007), 65(5),
                          CODEN: APFRAD; ISSN: 0003-4509
PUBLISHER:
                          Elsevier Masson SAS
DOCUMENT TYPE:
LANGUAGE:
                          English
    Per(6-iodo-6-desoxy) cyclodextrins are synthesis intermediates used in the design of the cation chelating per(3,6-anhydro)
     cyclodextrins. The modifications of the properties of these mols.
resulting from the nature of the halogen substituent and also the number of
     osidic building blocks were investigated by varying both factors, using 1H
     and 31P-NMR and EPR spectroscopies. These nearly water insol. mols.
     exhibits no complexing properties (for both ionic and apolar structures)
     but can be partially solubilized in micelles of detergent (SDS) and also
     in phospholipid vesicles. Dipolar connectivity (nOesy) NMR expts. show
     that they are embedded at the chain level of the micelles/vesicles,
     without any inclusion complex formation. Changing the number of glucose
     building blocks (6,7 or 8) or/and the nature of the halogen nuclei at the
     positions 6 strongly modify <u>cyclodextrin</u> affinities and membrane interactions. For instance the per(6-bromo-6-desoxy)-cyclomaltohexaose
     (ABR) and -cyclomalto-heptaose (BBR) exhibit a selective affinity for
     cobalt (apparent Ka of 2500 and 790 M-1, resp.). In terms of interactions
     with membranes, \alpha derivs. induce sterical hindrance at the
     phosphorus level while destructuring the chains. Other derivs. are
     located deeper and rigidify the most superficial part of the chain,
     suppressing the jump in membrane fluidity at transition temperature
                                THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                          28
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
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L11 ANSWER 3 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          2006:40820 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                          Cation complexing 2-0-alkylated, 3,6-anhydro-a-
                          cyclodextrins: the side-chain length governs
                          physicochemical properties and practical applications
                          Pailler, J. Y.; Gadelle, A.; Fauvelle, F.;
Dabouis, V.; Crouzier, D.; Debouzy, J. C.
Unite de Biophysique, CRSSA, La Tronche, 38702, Fr.
CORPORATE SOURCE:
                          Journal of Drug Delivery Science and Technology
                          (2005), 15(6), 419-426
                          CODEN: JDDSAL; ISSN: 1773-2247
PUBLISHER:
                          Editions de Sante
DOCUMENT TYPE:
LANGUAGE:
                          English
AB A series of chain-grafted per-3,6-anhydro-α- cyclodextrins
(ACD) were synthesized and their cation complexing properties studied by
     1H-NMR spectroscopy. Superficial tension measurements, 1H-NMR
     spectroscopy and phase diagrams showed that the properties of ACD were
     closely related to LogP, which also controlled their interactions with
     membranes. As a result, practical applications could be proposed and
     further perspectives suggested. Hence direct decontamination in liqs. may
     be possible for most amphiphilic derivs., since these amphiphilic mols.
     form gels or soaps. The most hydrophobic derivative realizes an insol.
     complex that can be used for depollution or cation determination in liqs.
                                THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                          41
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 4 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
                          2005:548189 CAPLUS <<LOGINID::20080328>>
ACCESSION NUMBER:
DOCUMENT NUMBER:
                          Hexakis (3,6-anhydro) tetrakis (2A,B,D,E-O-butyl)
                          cyclomalto hexaose as a promising biological cation
                          cryptant: Complexation and NMR study of interaction
                          with membranes
AUTHOR(S):
                          Pailler, J.-Y.; Gadelle, A.; Debouzy, J.-C.
CORPORATE SOURCE:
                          CRSSA, Unite de Biophysique, La Tronche, 38702, Fr.
SOURCE:
                          Journal of Drug Delivery Science and Technology
                          (2005), 15(3), 237-244
                          CODEN: JDDSAL
PUBLISHER:
                          Editions de Sante
DOCUMENT TYPE:
                          Journal
LANGUAGE:
   Per anhydro α- cyclodextrin exhibits in vivo and in vitro
     cation complexation properties, especially for heavy metal cations. In order to
     enhance the selectivity for toxic cations, several alkyl derivs. were
     prepared by substitution at the C-2 position. Among the series of
     3,6-anhydro-a- cyclodextrin derivs. (from hexakis (3,6-anhydro) hexakis (2A,B,C,D,E,F-O-methyl) cyclomaltohexaose (M36) to
     hexakıs (3,6-anhydro) tetrakis (2A,B,D,E-O-octyl) cyclomaltohexaose (036)
     alkyl derivs.), hexakis (3,6-anhydro) tetrakis (2A,B,D,E-O-butyl)
     cyclomaltohexaose (B36) was found to be of special interest. The
     properties of B36 in aqueous solution and in the presence of synthetic membranes
     were studied by mass spectroscopy, 31P, 2H and 1H-NMR spectroscopy, by
     surface plasmon resonance using BIAcore, and via superficial pressure
     measurements. It was found that B36 exhibits a special affinity for lead
     compared to other heavy toxic cations (mercury, cadmium, uranyl), but a
     negligible affinity for physiol. cations (sodium, calcium, potassium),
     i.e., a great selectivity. The surface-active properties of the soapy B36
     solution in water (with DMSO < 5%) were determined by surface tension
     measurements. In terms of solubility, B36 is very soluble in methanol (30 mM),
     less in ethanol (2 mM), while poorly soluble in water (500 \mu M). However,
     the use of a ternary solvent system (methanol, ethanol, water) allowed the
     formation of a true gel. This, related with its amphiphilic properties
     and possibilities for peculiar interactions with membranes are shown by
     31P and 2H-NMR spectroscopic studies.
                                THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                          3.3
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 5 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                         2005:510596 CAPLUS <<LOGINID::20080328>>
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DOCUMENT NUMBER:

144:89979

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High-resolution solid-state 13C NMR study of
                            per(3,6-anhydro)-α- cyclodextrin based
polymers and of their chromium complexes
                             Cadars, Sylvian; Foray, Marie-Francoise; Gadelle,
                             Andree; Gerbaud, Guillaume; Bardet, Michel
CORPORATE SOURCE:
                             Service de Chimie Inorganique et Biologique,
                             Departement de Recherche Fondamentale sur la Matiere
                             Condensee, CEA-Grenoble, Grenoble, F-38054, Fr.
                             Carbohydrate Polymers (2005), 61(1), 88-94
                             CODEN: CAPOD8; ISSN: 0144-8617
PUBLISHER:
                             Elsevier B.V.
DOCUMENT TYPE:
LANGUAGE:
                            English
    High-resolution solid-state 13C NMR was employed to characterize polymers
      made of per-3,6-anhydro-\alpha- cyclodextrins with 1,6-drisocyanatohexane used to bridge the macrocycles. These materials
      were designed because of their insoly, and their extractant properties due
     to the presence of the <u>cyclodextrin</u> rings. The properties of this new type of material appear very promising as potential extractant of
     different oxoanions. The properties of these materials to bind chromate
      or dichromate ions appear to be particularly attractive since the extraction of
      chromium is high and moreover there is no degradation of the polymers that can
      be further regenerated. These features rely mostly on qual. and quant.
      analyses of CP/MAS spectra. The studies of the NMR relaxation times, TCH,
      TipH and TiC for the starting polymers and its metal complexes allowed
      obtaining valuable insights concerning the mol. sites of interactions of
      the polymers with the oxoanions.
REFERENCE COUNT:
                           2.0
                                   THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS
                                   RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 6 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                             2005:78816 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                             Inclusion complexes of trivalent lutetium cations with
                             an acidic derivative of per(3,6-anhydro)-\alpha-
                             Cyclodextrin
Bonnet, Celia; Gadelle, Andree; Pecaut,
AUTHOR(S):
                             Jacques; Fries, Pascal H.; Delangle, Pascale
CORPORATE SOURCE:
                             Laboratoire de Reconnaissance Ionique, SCIB,
                             CEA/DSM/DRFMC, CEA-Grenoble, Grenoble, 38 054, Fr.
SOURCE:
                             Chemical Communications (Cambridge, United Kingdom)
                             (2005), (5), 625-627
                             CODEN: CHCOFS: ISSN: 1359-7345
PUBLISHER:
                            Royal Society of Chemistry
DOCUMENT TYPE:
                            Journal.
LANGUAGE:
                            English
     The <u>cyclodextrin</u> derivative (hexakis(2-0-carboxymethyl-3,6-anhydro)-\alpha- <u>cyclodextrin</u> (H6ACX)) forms mono- and bimetallic complexes with bu(III) in aqueous solution The x-ray structure of binuclear
      [Lu2(ACX)(H2O)2] is the 1st example of a lanthanide-cyclodextrin inclusion complex. The stability consts. of Lu-H6ACX complexes were determined
                            20
                                   THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                                   RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 7 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                            2004:83689 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                            Hydrolytic properties of per (3,6-anhydro,
                            2-O-carboxymethyl) alpha cyclodextrin complexes of Ce (III) and Eu (III): application to
                             soman (GD) degradation
AUTHOR(S):
                             Debouzy, J. C.; Gadelle, A.; Fauvelle, F.;
                             Testylier, G.
CORPORATE SOURCE:
                             CRSSA, La Tronche, Fr.
                             Bollettino Chimico Farmaceutico (2003), 142(3),
                             CODEN: BCFAAI; ISSN: 0006-6648
PUBLISHER:
                            Societa Editoriale Farmaceutica
DOCUMENT TYPE:
LANGUAGE:
                            English
    Per (3,6-anhydro-2-0-carboxymethyle) α- cyclodextrin
      ([ACX]) is a polydentate analog of EDTA a well-known cation chelating
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reagent. ACX exhibits strong affinities in vitro for uranyl, cobalt and
     also for lanthanides such as Europium and Cerium. The hydrolytic
     activities of ACX-Eu and ACX-Ce complex were directly tested on an
     organophosphorous compound: the neurotoxic Soman (GD), an inhibitor of
     acetylcholinesterase (ACHE from rat brain). It was found a three fold
     reduction of soman activity when measured in the presence of Ce-ACX complex.
     Conversely, Eu-ACX effect did not result in soman inhibition variation
     under physiol. conditions. It is suggested that, considering usual
     organometallic complex of cyclodextrin, such direct complexes would be of interest in the design of pseudo-enzyme systems for
     phosphoester hydrolysis.
REFERENCE COUNT:
                                  THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS
                           2.5
                                  RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 8 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                           2003:990981 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                           Per(3,6-anhydro) cyclodextrin derivatives,
their preparation, and their use for the separation or
                            fixation of anions based on manganese and chromium
INVENTOR(S):
                           Gadelle, Andree
Commissariat A L'energie Atomique, Fr.; Centre
PATENT ASSIGNEE (S):
                            National De La Recherche Scientifique Cnrs
SOURCE:
                            Fr. Demande, 42 pp.
                            CODEN: FRXXBL
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                           KIND
                                  DATE
                                                 APPLICATION NO.
                                                                           DATE
     FR 2840906
                                   20031219
                            A1
                                                                          20020612
     FR 2840906
                            В1
     WO 2003106507
                            A1
                                                WO 2003-FR1741
                                                                           20030611
          W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
               CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
               GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
               LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM,
               PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT,
               TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
          RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
               KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
               FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
               BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     AU 2003250357
                            A1
                                                AU 2003-250357
                                                                           20030611
     EP 1511774
                            A1
                                   20050309
                                                EP 2003-760007
                                                                           20030611
          R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
               IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
                                   20051117
                                                 JP 2004-513337
                                                                          20030611
     JP 2005534729
     US 2006014722
                            A1
                                                 US 2005-517582
                                                 FR 2002-7205
PRIORITY APPLN. INFO.:
                                                                       A 20020612
                                                                      W 20030611
                                                 WO 2003-FR1741
OTHER SOURCE(S):
                           MARPAT 140:52345
    Derivs. of per(3,6-anhydro) cyclodextrins having the general formulas (I) and (II) are prepared which can be used for the separation or
     fixation of chromate, dichromate and/or manganate anions from water or as
     a pharmaceutical complexing agent for humans. R1 in the general formulas
      I and II represents -OCONHR2, OH, OR3, SH, SR3, OCOR3, NH2, NHR3, NR3R4,
     CONH2, CONR3R4, CN, COOR3, OCH2COOH, or COOH, R3 and R2 represent an
     aliphatic, saturated or unsatd. group, R3 and R4 represent an aliphatic or aromatic
     hydrocarbon group which can be saturated or unsatd. and which can be
     substituted by halogen atoms or hetero atoms, such as O, S, and N, and n
     is 6, 7, or 8, or R1 represents the group OCONH(CR5R6)mNHCOOR7 with R5 and
     R6 being aliphatic saturated or unsatd, groups, and R7 represents glucosidic or
     maltosidic units of peranhydrocyclodextrin and m is a number from 1 to 20.
     Preferably, R1 of the per(3,6-anhydro) cyclodextrin derivative is -OCONHR2 with R2 being an Et or hexyl group and n being 6. The
     per(3,6-anhydro) <u>cyclodextrin</u> derivs. are prepared by reacting per(3, 6-anhydro) <u>cyclodextrins</u> having the general formulas (III) and (IV) with an isocyanate OCN-R2 or a diisocyanate OCN (CRSR6)mNCO.
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Polymers are obtained by reacting at least two per(3,6-anhydro)

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cyclodextrin derivs. having the general formulas III and IV with n
     and m being 6 and R5 and R6 being H. For the removal of anions from water
     the per(3,6-anhydro) cyclodextrin derivative or polymer is dissolved
     in an organic solvent immiscible with water.
                          6
                                 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 9 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
                           2003:940046 CAPLUS <<LOGINID::20080328>>
ACCESSION NUMBER:
DOCUMENT NUMBER:
                           In vitro cellular toxicity and in vitro lethality
                           studies of alkylated a-anhydro
                           cyclodextrins
Debouzy, J. S.; Gadelle, A.; Pailler, J. Y.;
Fusai, T.; Dabouis, V.; Pradines, B.; Fauvelle, F.;
AUTHOR(S):
                           Crouzier, D.
CORPORATE SOURCE:
                           CRSSA/BCM et Service d'Imagerie, La Tronche, 38702,
                           STP Pharma Sciences (2003), 13(3), 209-214
                           CODEN: STSSE5; ISSN: 1157-1489
                           Editions de Sante
DOCUMENT TYPE:
LANGUAGE:
                           English
    The overall toxicity of several per(3, 6-anhydro)-\alpha-
     cyclodextrins was studied both in vivo, in mice (mortality), and in vitro, in cells (VERO and CHO strains) and erythrocytes (hemolytic
     activity). It was found that mortality increased with the chain length,
     thus ranging from 0% (35 mM, saturated solution of per(3,6-anhydro)-\alpha-
               rin, A36) to a LD50 of 45-48 mM (per(2-0-methyl), M36)),
     and to 30% death at 10 mM (saturated per(2-0-Et, E36). A similar dependence
     of hemolytic activity on the chain length was also found, with the lowest
     HD50 observed for E36 and a negligible hemolysis observed for A36 and M36.
     Furthermore, cell toxicities observed on VERO and CHO cell cultures provided
     guite similar results. Finally, E36 was the only derivative able to interfere
     with the cell adhesiveness in plasmodium infected cells. It was suggested
     that the tensioactive properties of E36 are related both with this
     activity and with the overall toxicity of these derivs. Other chemical
     modifications were proposed to enhance the security range between toxicity
     and anti-adhesive activity.
                                 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                           39
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 10 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                           2003:102935 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                           139:129243
                           In vitro uranyl affinity of per(3,6-anhydro-2-o-
                           carboxymethyl)-α- cyclodextrin and
conditions required for in vivo application
                           Debouzy, J. C.; Gadelle, A.; Tymen, H.; Le Gall, B.; Millot, X.; Moretto, P.; Fauvelle, F.; Le
AUTHOR(S):
                           Peoc'H, M.; Dabouis, V.; Martel, B.
                           UMR 5046, CEA/DRFMC/SCIB/FI, Grenoble, F38054, Fr.
CORPORATE SOURCE:
SOURCE:
                           Annales Pharmaceutiques Françaises (2003), 61(1),
                           CODEN: APFRAD; ISSN: 0003-4509
PUBLISHER:
                           Masson Editeur
DOCUMENT TYPE:
                           Journal
LANGUAGE:
                           French
AB Per(3.6-anhydro-2-0-carboxymethyle)-a-cyclodextrin ([I]) is a polydentate analog of EDTA, a well-known cation chelating reagent. I
     exhibits strong affinities in vitro for lanthanides, cobalt and also for
     uranyl cations. A 1:1 stoichiometry and a high affinity for uranyl
     (6<logK<7) were found in vitro. I is not hemolytic and exhibits no lethal properties in mice (LD50 42 mM). In vivo injection at supralethal amts.
     of uranyl complex of I prevents immediate death in mice, while it is
     unable to protect against later death. Pharmacokinetic studies show that
     a dissociation of the complex occurs, leading to the release of free uranyl.
     Complexation assays of I, Co nitrate and Pb nitrate, using
     cyclodextrin-functionalized polyester fabrics were also carried
                                 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS
                                  RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
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L11 ANSWER 11 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
                          2003:68010 CAPLUS <<LOGINID::20080328>>
ACCESSION NUMBER:
DOCUMENT NUMBER:
                          First evaluation of per(3,6-anhydro,2-0-carboxymethyl) -
                          α- cyclodextrin for biological
                          decontamination of cobalt
AUTHOR(S):
                          Debouzy, J. C.; Tymen, H.; Le Gall, B.; Fauvelle, F.;
                          Martel, B.; Gadelle, T.; Gadelle, A.
Unite de Biophysique et Service de Biospectrometrie,
CORPORATE SOURCE:
                          CRSSA, La Tronche, 38702, Fr.
                          S.T.P. Pharma Sciences (2002), 12(6), 397-402
                          CODEN: STSSE5; ISSN: 1157-1489
PUBLISHER:
                          Editions de Sante
DOCUMENT TYPE:
LANGUAGE:
AB Per (3,6-anhydro-2-0-carboxymethyl)-a- cyclodextrin (ACX)
     is a polydentate analog of EDTA, a known cation chelating reagent. ACX
     exhibits strong affinities in vitro for lanthanids, uranyle and especially for
     Co. The possible application of ACX for Co decontamination was tested in
     an aqueous solution and incorporated in agarose gel on human skin (in Franz's
     diffusion chambers) and living rats. In comparison with EDTA and DTPA,
     skin decontamination by ACX was better when it was incorporated in a gel
     and similar after several skin washing cycles. Several ACX-loaded tissues
     (viscose and polyester) were also assayed on the same model and showed an
     increased fixation of Co by ACX-loaded viscose, whereas this was not observed
     with polyester.
REFERENCE COUNT:
                          2.3
                                THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 12 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          2002:923095 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                          Amphiphilic per (3,6-anhydro, 2-0-ethyl)-a-
                          cyclodextrin: the first step towards
                          self-gelifying cation cryptants?
                          Debouzy, J. C.; Gadelle, A.; Fauvelle, F.;
Pailler, J. Y.; Brasme, B.; Dabouis, V.; Aous, S.;
AUTHOR(S):
                          Fusai. T.
CORPORATE SOURCE:
                          Unite de Biophysique et Service de Biospectrometrie,
                          CRSSA, La Tronche, 38702, Fr.
SOURCE:
                          S.T.P. Pharma Sciences (2002), 12(5), 267-273
                          CODEN: STSSE5; ISSN: 1157-1489
PUBLISHER:
                          Editions de Sante
DOCUMENT TYPE:
                          Journal.
LANGUAGE:
                          English
    The properties of per(3,6-anhydro, 2-0-ethyl)-a- cyclodextrin
     (3,6-CDE) in solution and in the presence of synthetic membranes were studied
     by thin layer chromatog., mass, 31P-, 2H- and 1H-NMR spectroscopies, and
     superficial pressure measurements. It was found that 3,6-CDE exhibits a
     good affinity for Co2+, Hg2+, Sr2+, Pb2+ and Na+. Besides, ROESY expts.
     showed that two different conformations of 3,6-CDE were simultaneously
     present during slow exchange. The tensioactive properties of the soapy
     solution of 3,6-CDE in water/ethanol were shown by superficial tension (ST)
     measurements. Moreover, 31P-NMR showed an increase of the superficial
     fluidity of phospholipid dispersions, above the transition temperature in the presence of 3,6-CDE. Furthermore, no detergent effect was observed in the
     presence of small unilamellar vesicles of lecithin, membrane destructions
     being only observed after several days, or when 3,6-CDE and phospholipids
     were co-sonicated. These results lead to the discussion of the biol.
     availability of 3,6-CDE as a wound decontaminant, further chemical
     modifications being also suggested.
                         41
                                THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 13 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          2002:514937 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                          Biocompatible gels comprising peranhydrodextrins
                          useful for decontaminating wounds contaminated by
                          heavy metals such as lead
INVENTOR(S):
                          Baudin, Cecile; Perly, Denis; Gadelle, Andree
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; Debouzy, Jean Claude; Fauvelle, Florence
PATENT ASSIGNEE(S):
                          Commissariat a l'Energie Atomique, Fr.
                          Fr. Demande, 14 pp.
                          CODEN: FRXXBL
DOCUMENT TYPE:
                          Patent
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                                 DATE
                                              APPLICATION NO.
                                                                       DATE
     FR 2814748
                           A1
PRIORITY APPLN. INFO.:
OTHER SOURCE(S):
                         MARPAT 137:52362
AB Biocompatible gels comprising peranhydrodextrins, a gelling agent, and
     water are useful for decontaminating wounds contaminated by heavy metals
     such as lead. A gel contained permethyl-perhydro-α-
     cyclodextrin 20, agarose 3 g/L.
L11 ANSWER 14 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          2002:484484 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                          Acidic Derivative of Per(3,6-anhydro)-a-
                          cyclodextrin: Preparation and a First
                          Evaluation of Its Affinity for Lanthanides by 1H NMR
AUTHOR(S):
                          Fauvelle, F.; Gadelle, A.; Pailler, Y.; Aous, S.; Debouzy, J. C.
CORPORATE SOURCE:
                          Laboratoire de Biophysique, CRSSA, La Tronche, 38702,
SOURCE:
                          Journal of Inclusion Phenomena and Macrocyclic
                          Chemistry (2002), 42(3-4), 203-207
                          CODEN: JIPCF5; ISSN: 1388-3127
                          Kluwer Academic Publishers
PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
                          English
OTHER SOURCE(S):
                          CASREACT 137:353231
   We report on the first synthesis of hexakis(2-0-carboxymethyl-3,6-anhydro)-
     a- cyclodextrin, an acidic derivative of per(3,6-anhydro)-
    a- cyclodextrin. Preliminary qual. tests showed that this new compound would have greater affinity for lanthanides, cobalt and uranyl
     cations, than for sodium, potassium and calcium physiol. ions.
REFERENCE COUNT:
                                THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 15 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          2001:730839 CAPLUS <<LOGINID::20080328>>
                          135:290396
DOCUMENT NUMBER:
                          Per(3,6-anhydro) cyclodextrin derivatives,
preparation and use thereof for separating ions
                          Gadelle, Andree; Fauvelle, Florence;
INVENTOR(S):
                          Debouzy, Jean-Claude
PATENT ASSIGNEE (S):
                          Commissariat a l'Energie Atomique, Fr.; Centre
                          National de la Recherche Scientifique (CNRS)
                          PCT Int. Appl., 32 pp.
SOURCE:
                          CODEN: PIXXD2
DOCUMENT TYPE:
                          Patent
LANGUAGE:
                          French
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
    PATENT NO.
                          KIND
                                 DATE
                                              APPLICATION NO.
                                                                       DATE
     WO 2001072849
                          A1
                                 20011004
                                              WO 2001-FR923
                                                                       20010327
         W: US
         RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE, TR
     FR 2807044
                           A1
                                              FR 2000-3899
                                                                       20000328
    FR 2807044
                           B1
                           A1
                                              EP 2001-919576
                                                                       20010327
                           В1
                                 20041110
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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AT 282048
                                             AT 2001-919576
                                             ES 2001-919576
     ES 2231469
                                20050516
     US 2002137923
                                             US 2001-926637
                          A1
                                                                 A 20000328
PRIORITY APPLN. INFO.:
                                             FR 2000-3899
                                             WO 2001-FR923
                                                                 W 20010327
OTHER SOURCE(S):
                         MARPAT 135:290396
    The invention concerns per(3,6-anhydro)cyclodextrin derivs.,
     their preparation and their use for separating polluting ions, for example, for
     human decontamination. The derivs, bear axially or equatorially
     substituted group R1 on positions 2 where one R1 at least represents the
     -OCH2COOH group and the other R1's, identical or different, correspond to
     one of the formulas: OH, OR2, SH, SR2, OCOR2, NH2, NHR2, NR2R3, CONH2,
     CONHR2, CONR2R3, CN, COOR2, COOH and R2, wherein: R2 and R3, identical or
     different, represent a saturated or unsatd. hydrocarbon, aliphatic or aromatic
     group, capable of comprising one several heteroatoms selected among 0, 8
     and N; and n is equal to 6, 7 or 8. Thus, heating 1 g
     hexakis(3,6-anhydro)cyclomaltohexaose for 2 h at 120°, adding 10 mL
     DMSO and 10 mL a 2N NaH DMSO solution, mixing under Ar for 3 h at room temperature,
     combining the resulting blue-gray solution with 1.6 g Na monochloroacetate,
     mixing at room temperature for 24 h and working up gave a hexakis (3,6-anhydro-2-
     O-carboxymethyl)cyclomaltohexaose which formed easily complexes with aqueous
     solution containing Lu3+, La3+, Dy3+, Eu3+ and Co2+ ions.
                               THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                         4
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 16 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                         2001:341369 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                          1H-NMR study of heavy metals complexation with
                         hexakis (3,6-anhydro) tetrakis (2A,B,D,E-0-
                          octyl)cyclomaltohexaose (oct)
                         Debouzy, J. C.; Gadelle, A.; Fauvelle, F.;
Nardin, R.; Aous, S.; Lhoste, F.; Pailler, Y.
AUTHOR(S):
CORPORATE SOURCE:
                          CRSSA, Biological and molecular biophysics Lab., La
                          Tronche, Fr.
                          Bollettino Chimico Farmaceutico (2001), 140(1), 9-14
                         CODEN: BCFAAI; ISSN: 0006-6648
PUBLISHER:
                          Societa Editoriale Farmaceutica
DOCUMENT TYPE:
                          Journal.
LANGUAGE:
                         English
     The selection of cations bound by hexakis (3,6-anhydro) tetrakis
     (2A, B, D, E-O-octyl) cyclomaltohexaose (OCT) was performed by thin layer
     chromatog. The 3 cations selected, UO22+, Pb2+ and Hg2+, were then
     studied by 1H-NMR. A 2:1 OCT/cation stoichiometry was identified in the
     cases of UO22+ and Pb2+. While UO22+ binding (log K around 6) followed a
     fast exchange kinetics, a slow or intermediate complexation was observed with
     Pb2+ (log K-5.6) and Pb2+, resp. In the latter case, because of the the
     poor solubility of Hg2+, neither a stoichiometry nor an estimation of the affinity
     constant could be proposed.
                         19
                               THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS
                               RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 17 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          2000:729064 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                          134:17643
                          2-0-substituted-3,6-per-anhydro-a-
                          cyclodextrin as potential biocompatible agents
                          for the selective complexation of heavy metal ions
                          with special attention to lead
AUTHOR(S):
                          Baudin, Cecile; Pean, Christophe; Pellizzari, Bruno;
                          Gadelle, Andree; Fauvelle, Florence; Debouzy,
Jean-Claude; Dalbiez, Jean-Pierre; Perly, Bruno
CORPORATE SOURCE:
                          CEA, DRECAM/SCM, CEN de Saclay, Gif sur Yvette,
                          F-91191, Fr.
                          Journal of Inclusion Phenomena and Macrocyclic
                          Chemistry (2000), 38(1-4), 287-296
                         CODEN: JIPCF5
                         Kluwer Academic Publishers
DOCUMENT TYPE:
LANGUAGE:
                         English
AB We report on the synthesis, characterization and ionic complexation
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properties of hexakis (2-0-acetyl-3,6-anhydro) cyclomaltohexaose and
     hexakıs (2-0-methyl-3,6-anhydro) cyclomaltohexaose using thin-layer
     chromatog, and NMR spectroscopy. The selectivity towards cations depends
     on chemical modification of the hydroxyl groups and a very high specificity
     can be obtained in the case of lead for methylated derivs.
REFERENCE COUNT:
                                THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 18 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                           2000:311144 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                           132:339914
                           Cation complexation properties of hexakis(2-0-methyl-
                           3,6-anhydro)-a- cyclodextrin: A 1H NMR
AUTHOR(S):
                          Fauvelle, F.; Gadelle, A.; Debouzy, J. C.; Baudin, C.; Perly, B.
CORPORATE SOURCE:
                          CRSSA, laboratoire de Biophysique, La Tronche, 38702,
                           Supramolecular Chemistry (2000), 11(3), 233-237
                           CODEN: SCHEER; ISSN: 1061-0278
                           Gordon & Breach Science Publishers
DOCUMENT TYPE:
LANGUAGE:
    The affinity of hexakis(2-0-methyl-3,6-anhydro)-α-
     cyclodextrin (3,6-\alpha-CDM) for Ba2+, Pb2+, Ca2+ and Sr2+ has been tested by 1H NMR. 3,6-\alpha-CDM forms strong complexes in water
     with Pb2+ and Ba2+. The comparison with the parent hexakis(3,6-anhydro)-
     \alpha- cyclodextrin bearing hydroxyl groups instead of methoxy groups reveals that the O-CH3 substitution significantly improves the
     anhydro-cyclodextrin selectivity.
RENCE COUNT: 13 THERE A
REFERENCE COUNT:
                                 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 19 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          2000:56447 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                           Comparative cation chelating properties of
                           per(3,6-anhydro)- and per(3,6-anhydro 2-0 Me) \alpha-
                           cyclodextrins
Debouzy, J. C.; Fauvelle, F.; Gadelle, A.;
AUTHOR(S):
                           Dabouis, V.; Perrin, A.; Brasme, B.; Peinequin, A.;
                           Perly. B.
CORPORATE SOURCE:
                           CRSSA/Biophysics, La Tronche, 38702, Fr.
SOURCE:
                           Proceedings of the International Symposium on
                           Cyclodextrins, 9th, Santiago de Comostela, Spain, May
                           31-June 3, 1998 (1999), Meeting Date 1998, 105-108.
                           Editor(s): Labandeira, J. J. Torres; Vila-Jato, J. L.
                           Kluwer Academic Publishers: Dordrecht, Neth.
                           CODEN: 68NHAE
DOCUMENT TYPE:
                          Conference
LANGUAGE:
                          English
     The cation chelating properties of per(3,6 anhydro)-\alpha-
     cyclodextrin, [A36] and of per(3,6 anhydro, 2-0 Me)-a-
      cyclodextrin, [A36M] were studied by mass and NMR spectroscopy.
     A36 forms 1:1 complexes with lead (K = 2500 M-1), and also with Sr and K
     with a fast exchange rate kinetics. However, the formation of A36-Pb
     complex results in a dramatic enhancement of the hemolytic properties.
     Permethylation at the position 2 (A36M) confers an extreme affinity for
     Ba2+, Pb2+, Sr2+ and Ca2+ following a slow rate exchange process and a 1:1
     stoichiometry. A weak 1:1 A36M-K complex is also found with a fast
     exchange rate. In contrast to A36, A36M complexes showed no hemolytic
     properties. An agarose gel of A36M was successful in the decontamination
     of wounds polluted with lead or strontium ions on rats.
REFERENCE COUNT:
                                 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 20 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          2000:56439 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                           132:222741
                           Mono-6-tosyl-β- cyclodextrin:
                          preparation, hydrolysis and self-inclusion studies in
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aqueous solution

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AUTHOR(S):
                          Djedaini-Pilard, F.; Gosnat, M.; Steinbruckner, S.;
                           Dalbiez, J. P.; Crini, G.; Perly, B.; Gadelle,
CORPORATE SOURCE:
                           DRECAM/SCM, CEA-Saclay, Gif sur Yvette, F-91191, Fr.
                           Proceedings of the International Symposium on
                           Cyclodextrins, 9th, Santiago de Comostela, Spain, May
                           31-June 3, 1998 (1999), Meeting Date 1998, 73-76.
                           Editor(s): Labandeira, J. J. Torres; Vila-Jato, J. L.
                           Kluwer Academic Publishers: Dordrecht, Neth.
                           CODEN: 68NHAE
DOCUMENT TYPE:
LANGUAGE:
   We show here that the kinetics of the reaction of tosylation in aqueous solution
     strongly depends upon the effective pH. In alkaline aqueous solution, although the
     reaction is very fast and can yield up to 35% of the title compound, it is
     competing with hydrolysis of the mono-6-tosyl-6-deoxy-\beta-
     cyclodextrin (1). A complete NMR study has demonstrated that this
     product is hydrolyzed in aqueous solution at pH > 6 and that acidification of the
     reaction medium can quench this process. Investigations of the structure
     of pure 1 in aqueous solution are presented showing that a strong intramol.
     self-inclusion complex is formed. Dedicated two dimensional NMR expts.
     are used in conjunction with competition with external quests to evidence
     and estimate the strength of the auto-inclusion complex.
REFERENCE COUNT:
                                 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 21 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
                           1999:535329 CAPLUS <<LOGINID::20080328>>
ACCESSION NUMBER:
DOCUMENT NUMBER:
                           132:88121
                           Interaction of per(3,6-anhydro)-\alpha-
                           cyclodextrin (a36CD) and
lead-a36CD complex with biological systems
                          Debouzy, J. C.; Fauvelle, F.; Gadelle, A.;
Baudin, C.; Richard, M.; Perly, B.; Chouteau, F.;
AUTHOR(S):
                           Joets, J.; Tazz, J. J.; Daveloose, D.
CORPORATE SOURCE:
                           CRSSA, Laboratoire RMN, Tronche, 38702, Fr.
SOURCE:
                           Bollettino Chimico Farmaceutico (1998), 137(5),
                           144-151
                           CODEN: BCFAAI; ISSN: 0006-6648
PUBLISHER:
                          Societa Editoriale Farmaceutica
DOCUMENT TYPE:
LANGUAGE:
                          English
    The interactions of per(3,6 anhydro)-a- cyclodextrin
     (α36CD) and of lead-α36CD complex with biol. systems were
     tested by NMR, ESR and electronic microscopy using erythrocytes and model
     membranes. It was found that the hemolytic activity of \alpha 36\text{CD} alone
     was seven fold lower than that of natural \alpha- cyclodextrin (evaluated by the concentration inducing 50% hemolysis, DH50-35 mM). Conversely,
     the formation of the complex resulted in an increase of hemolytic
     properties, with DH50 of 1 mM. The mechanism proposed was an increased
     membrane diffusion by endocytosis of the complex, leading to higher amts.
     of intracellular lead.
REFERENCE COUNT:
                                 THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 22 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
                           1999:68297 CAPLUS <<LOGINID::20080328>>
ACCESSION NUMBER:
DOCUMENT NUMBER:
                           The cation complexation properties of
                           per-3,6-anhydro-\alpha and \beta-
                           cyclodextrins studied by thin layer
                           chromatography and 1H NMR
AUTHOR(S):
                           Fauvelle, F.; Gadelle, A.; Debouzy, J. C.;
                           Perly, B.
CORPORATE SOURCE:
                           CRSSA, Biophysique, La Tronche, 38702, Fr.
SOURCE:
                           Molecular Recognition and Inclusion, Proceedings of
                           the International Symposium on Molecular Recognition
                           and Inclusion, 9th, Lyon, Sept. 7-12, 1996 (1998),
Meeting Date 1996, 325-328. Editor(s): Coleman,
                           Annette W. Kluwer: Dordrecht, Neth.
                           CODEN: 67FSAY
DOCUMENT TYPE:
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LANGUAGE:
                          English
AB A step scale affinity of cations for per-3,6-anhydro-α-
     cyclodextrin (3,6-aCD) can be deduced from NMR binding
     constant determination which is in agreement with TLC results: Pb2+ » Sr2+ >
     K+ > Cs+ > NH4. The other ions tested, like Na+ and Ca2+, did not induce
     any observable spectral modifications on the NMR time-scale. The
     3,6-αCD mol. is then selective for Pb2+. Conversely, 3,6-βCD
     has poor cation binding properties: only K+ and Cs+ are complexed. The
     weakness of the binding consts. and the absence of selectivity are not in
     favor of a biol. use.
REFERENCE COUNT:
                                 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 23 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          1999:68293 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                          NMR study of per(3,6-anhydro)-\alpha-
                          cyclodextrin as a potential agent for the biological decontamination of lead as evidenced by NMR
                          spectroscopy
                          Debouzy, J. C.; Fauvelle, F.; Gadelle, A.;
                          Perly, B.; Baudin, C.
CORPORATE SOURCE:
                          CRSSA, U.Biophysique, La Tronche, 38702, Fr.
                          Molecular Recognition and Inclusion, Proceedings of
                          the International Symposium on Molecular Recognition
                          and Inclusion, 9th, Lyon, Sept. 7-12, 1996 (1998), Meeting Date 1996, 309-312. Editor(s): Coleman,
                          Annette W. Kluwer: Dordrecht, Neth.
                          CODEN: 67FSAY
DOCUMENT TYPE:
                          Conference
LANGUAGE:
                          English
     The ability of per(3,6-anhydro)-\alpha- cyclodextrin (A36CD) to capture lead from a preformed glutathione-lead complex was investigated by
     NMR spectroscopy. This strongly depends on the nature and pH of the
     buffer used in the competition expts. It was found that an almost
     complete removal of lead can be achieved at pH 5.5, especially when lead nitrate
     is used. The capture also strongly depends on the nature of the lead
     species as well as of the counter ion present in the medium. These
     observations imply that decontamination of lead by this process will be
     optimal under acidic conditions.
                          8
                                THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 24 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          1999:8034 CAPLUS << LOGINID::20080328>>
DOCUMENT NUMBER:
                          Method for fixing or separating ions such as lead by
                          using per(3,6-anhydro)cyclodextrin
                          derivatives
INVENTOR(S):
                          Baudin, Cecile; Perly, Bruno; Gadelle, Andree
                          ; Debouzy, Jean-Claude; Fauvelle, Florence
PATENT ASSIGNEE(S):
                          Commissariat a l'Energie Atomique, Fr.
                          PCT Int. Appl., 30 pp.
SOURCE:
                          CODEN: PIXXD2
DOCUMENT TYPE:
                          Patent
LANGUAGE:
                          French
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                          KIND
                                 DATE
                                              APPLICATION NO.
                                                                       DATE
     WO 9856829
                           A1
                                  19981217
                                              WO 1998-FR1235
                                                                       19980612
         W: AU, HU, JP, RU, US
         RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE
                           A1
                                              FR 1997-7339
                                                                       19970613
     ZA 9805079
                           A
                                              ZA 1998-5079
                                                                       19980611
                           Α
                                              AU 1998-82181
                                                                       19980612
     AU 752287
                           B2
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EP 1998-932194

EP 991670

EP 991670

A1

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R: CH, DE, GB, IT, LI, NL, SE
     HU 2000002298 A2 20001128
HU 2000002298 A3 20030528
                                               HU 2000-2298
                                                                        19980612
                                  20020205
                                                JP 1999-501800
                                                                         19980612
     US 6544964
                                  20030408
                                               US 2000-445818
PRIORITY APPLN. INFO.:
                                               FR 1997-7339
                                                                     A 19970613
                                               WO 1998-FR1235
                                                                     W 19980612
OTHER SOURCE(S):
                          MARPAT 130:71569
AB A method for fixing or separating ions, in particular of lead by using
     per(3,6-anhydro)cyclodextrin derivs. consists in contacting the medium containing the ions to be fixed or separated, with the derivative Preferably,
     for fixing lead hexakis(3,6-anhydro-2-0-methyl)cyclomaltohexaose (I) is
     used. The complexation will eliminate the environmental lead pollution.
     Thus, I was prepared by the methylation of hexakis(3,6-
     anhydro)cyclomaltohexaose with MeI in the presence of NaH in DMF solution I
     was then treated with Pb(NO3)2 to give the complex which was characterized
     by spectral methods. I is useful for the decontamination of lead.
REFERENCE COUNT:
                                 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 25 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                           1997:786657 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                           128:16383
                           Mechanism of \alpha- cyclodextrin induced hemolysis. 2. A study of the factors controlling the
                           association with serine-, ethanolamine-, and
                           choline-phospholipids
AUTHOR(S):
                           Debouzy, J. C.; Fauvelle, F.; Crouzy, S.; Chapron, Y.;
                           Goschl, M.; Gadelle, A.
Unite de Biophysique, CRSSA, La Tronche, 38702, Fr.
CORPORATE SOURCE:
                           Journal of Pharmaceutical Sciences (1998), 87(1),
                           59-66
                           CODEN: JPMSAE; ISSN: 0022-3549
PUBLISHER:
                           American Chemical Society
DOCUMENT TYPE:
LANGUAGE:
                           English
   A NMR spectroscopy and mol. modeling study of the interaction between
     α- cyclodextrin (α-CD) and phospholipids with serine, ethanolamine, or choline headgroups was based on 31P and 1H NMR
     measurements on small unilamellar vesicles (SUV), multilamellar vesicles
     (MLV), and aqueous suspensions of lipids using a direct complex preparation with
     α-CD. Mol. dynamics computer simulations were used to investigate
     the trajectory of \alpha-CD in the vicinity of a membrane surface and the
     influence of the charge and dipole moment of the phospholipid headgroups.
     These factors of charge and orientation of dipole moment seemed to play a
     key role in the interaction of phospholipids with \alpha\text{-CD} and reflected
     very well the exptl. observed selectivity of the approach of \alpha-CD to
     phospholipid. However, with this approach, there is no evidence for the
     formation of a complex with the phospholipid headgroup (except for
     phosphatidylinositol) that results from electrostatic forces. Rather,
     after a possible extraction of the lipid from the membrane, a classical
     inclusion of the sn-2 chain in the cavity of \alpha\text{-CD} occurs. This step
     depends on the alkyl chain length and saturation state of the lipids as well as
     on their organization (i.e., as vesicles or dispersions). Possible chemical
     modifications of the \alpha-\text{CD} mol. to control the hemolytic properties
     of \alpha-CD are discussed.
L11 ANSWER 26 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                           1997:697961 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                           A mild one-step selective conversion of primary
                           hydroxyl groups into azides in mono- and
                           oligosaccharides
AUTHOR(S):
                           Luis Jimenez, Jose Luis; Garcia Fernandez, Jose
                           Manuel; Gadelle, Andree; Defaye, Jacques
CSIC and Universidad de Sevilla, Instituto de
CORPORATE SOURCE:
                           Investigaciones Quimicas, Seville, E-41092, Spain
                           Carbohydrate Research (1997), 303(3), 367-372
                           CODEN: CRBRAT; ISSN: 0008-6215
PUBLISHER:
                           Elsevier
```

DOCUMENT TYPE: LANGUAGE:

English

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OTHER SOURCE(S):
                         CASREACT 127:359022
AB The direct azidation reaction of several monosaccharide Me
     glycopyranosides, sucrose, a,a-trehalose, cyclomaltohexaose
     and cyclomaltoheptaose with sodium azide in the presence of
     triphenylphosphine-carbon tetrabromide is reported. The optimal reaction
     conditions require pre-formation of the reactive species before addition of
     the sugar substrate. Formation of the primary azidodeoxy compound is
     accompanied by simultaneous formation of the corresponding primary
     bromodeoxy and 3,6-anhydro derivs. in the glycopyranoside series,
     former being transformed in situ into the azide by quenching of the
     reaction mixture with methanol before increasing the temperature Interestingly,
     good selectivity towards the primary C-6 position of the glucopyranosyl
     molety as compared to the fructofuranosyl one was observed in the case of
     sucrose, advantage of which has been taken in an improved preparation of
     2,3,4,1',3',4',6'-hepta-O-acetyl-6-azido-6-deoxysucrose (45% yield from
     sucrose). Sodium or lithium azide reagents were found equally effective.
     The azide functionality could be reduced without previous purification and the
     resulting amino sugar isolated by cation-exchange column chromatog., as
     illustrated for the preparation of 61-amino-61-deoxycyclomaltoheptaose.
REFERENCE COUNT:
                          23
                                THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 27 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          1997:606040 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                          The hemolytic properties of chemically modified
                          cyclodextrins
Bost, Mireille; Laine, Valerie; Pilard, Florence;
AUTHOR(S):
                                le, Andree; Defaye, Jacques; Perly, Bruno
                          Laboratoire d'Hematologie, Centre Hospitalier
CORPORATE SOURCE:
                          Universitaire de Grenoble, Grenoble, F-38043, Fr.
                          Journal of Inclusion Phenomena and Molecular
                          Recognition in Chemistry (1997), 29(1), 57-63
                          CODEN: JIMCEN; ISSN: 0923-0750
PUBLISHER:
                          Kluwer
DOCUMENT TYPE:
LANGUAGE:
                          English
    The hemolytic properties of natural cyclodextrins, especially of the more common cyclomaltoheptaose entity, severely hamper their potential use
     as carriers in pharmaceutical applications where parenteral administration
     is concerned. A systematic investigation on the role of chemical
     modifications with regard to the hemolytic character was carried out
     involving C-6 branched neutral, anionic, cationic and amphoteric derivs.
     From these data, conclusions have been drawn about the charge and the
     geometry of the modification: (1) substitution at primary hydroxyl groups
     usually decreases the hemolytic character and the geometry of the
     substituent affects the hemolytic property; (2) introduction of an amino
     group, resulting in a pos. charge at physicl. pH, decreases the hemolytic
     character; (3) neg. charges are comparatively less effective in reducing
     the hemolytic character; (4) zwitterionic groups seem to enhance the
     hemolytic character of the cyclodextrin mol.

PRINCE COUNT: 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS
                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L11 ANSWER 28 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          1997:553822 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                           127:190980
                          Substituted derivatives of per(3,6-anhydro)
                          cyclodextrins, process for their preparation and their uses for TLC separation of cations
INVENTOR(S):
                          Baudin, Cecile; Perly, Bruno; Gadelle, Andree
PATENT ASSIGNEE(S):
                          Commissariat a l'Energie Atomique, Fr.
SOURCE:
                          Eur. Pat. Appl., 6 pp.
                          CODEN: EPXXDW
DOCUMENT TYPE:
                          Patent
LANGUAGE:
                          French
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                                 DATE
                                              APPLICATION NO.
                                                                      DATE
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EP 787744

A1

19970806

EP 1997-400197

19970128

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EP 787744
                           В1
                                 20010613
         R: CH, DE, GB, IT, LI, NL, SE
     FR 2744124
                          A1
                                 19970801
                                              FR 1996-1073
                                                                      19960130
                                 19980306
                                              US 1996-773001
                                                                      19961223
     AU 9712303
                           Α
                                 19970807
                                              AU 1997-12303
                                                                      19970123
     AU 707604
                           В2
                                 19990715
     ZA 9700689
                          Α
                                              ZA 1997-689
                           Α
                                  19970812
     HU 9700280
                                 19971229
                                              HU 1997-280
                                                                      19970129
                                 20010129
PRIORITY APPLN. INFO.:
                                              FR 1996-1073
                                                                   A 19960130
                         MARPAT 127:190980
AB Per(3,6-anhydro)-(\alpha-, \beta-, and \gamma)- cyclodextrins, substituted at the 2' position with R (R = OH, OR1, SR1, OCOR1NH2, amine,
     amide, CONH2, CO2R1, OSO2R1, N3; R1 - H, alkyl, aryl, heterocycle) were
     prepared and used for TLC separation of cations. Thus, hexakis(3,6-anhydro-2-0-
     acetyl) cyclomaltohexaose was prepared and used for separation of cations, such as
     K+ and Cs+, by TLC .
L11 ANSWER 29 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          1996:178584 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                          Sensing effects for bioapplications in
                          electroconducting conjugated polymers
                          Bidan, Gerard; Gadelle, Andree; Teoule, Robert; Vieil, Eric
AUTHOR(S):
CORPORATE SOURCE:
                          Departement de Recherche Fondamentale sur la Matiere
                          Condensee, Centre d'Etudes Nucleaires de Grenoble,
                          Grenoble, F-38054, Fr.
                          Sensors and Materials (1996), 8(3), 179-84
SOURCE:
                          CODEN: SENMER; ISSN: 0914-4935
                          Scientific Publishing Division of MYU K.K.
DOCUMENT TYPE:
                          Journal
LANGUAGE:
                          English
AB Straightforward and easy electrodeposition of electroconducting conjugated
     polymers (ECPs) and their functionalization either by entrapment of anions
     or by covalent grafting make these materials attractive candidates for
     fabrication of a sensitive layer at the surface of an electrode. This
     approach is exemplified in a NO2--sensitive poly(N-methylpyrrole) layer,
     single-stranded DNA-derivatized polypyrrole film and a reservoir electrode
     based on a polypyrrole with host \beta- cyclodextrins.
L11 ANSWER 30 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                          1995:921924 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                          Method for solubilizing antitumor agents from the
                          taxol family in an aqueous medium, and branched
                          cyclodextrins therefor
                          Defaye, Jacques; Perli, Bruno; Gadelle, Andree
; Descamps, Valerie; Coste, Sarguet Annie
INVENTOR(S):
PATENT ASSIGNEE(S):
                          Commissariat a l'Energie Atomique, Fr.; Centre
                          National de la Recherche Scientifique
                          PCT Int. Appl., 29 pp.
                          CODEN: PIXXD2
DOCUMENT TYPE:
                          Patent
LANGUAGE:
                          French
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                          KIND
                                 DATE
                                              APPLICATION NO.
                                                                      DATE
     WO 9519994
                                  19950727
                          A1
                                              WO 1995-FR75
                                                                      19950124
         W: JP, US
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                                 19950728
                           A1
                                              FR 1994-778
                           В1
                                 19960405
PRIORITY APPLN. INFO.:
                                              FR 1994-778
                                                                   A 19940125
                         MARPAT 123:322100
AB According to the method, the antitumor agents of the taxol family were
```

solubilized by combining them with a branched cyclodextrin (I; n - 6-8; R1 - OH, SR2; R2 - α-maltosyl, β-maltosyl group).

L11 ANSWER 31 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN

1995:694615 CAPLUS <<LOGINID::20080328>>

DOCUMENT NUMBER: 124:9153

Inclusion and solubilization properties of

 $6-8-glycosyl-6-thio derivatives of <math>\beta$ -

Laine, Valerie; Coste-Sarguet, Annie; Gadelle, AUTHOR(S):

ndree; Defaye, Jacques; Perly, Bruno;

Djedaini-Pilard, Florence

CNRS, Centre d'Etudes de Grenoble, Grenoble, F-38054, CORPORATE SOURCE:

Journal of the Chemical Society, Perkin Transactions

2: Physical Organic Chemistry (1995), (7), 1479-87 CODEN: JCPKBH; ISSN: 0300-9580

Royal Society of Chemistry

DOCUMENT TYPE: LANGUAGE: English

CASREACT 124:9153

The synthesis and physico-chemical properties of branched β -

cyclodextrins substituted by one or seven thioglycoside units at the primary hydroxy side are described. The solubilities in water of these compds, are strongly increased compared with the parent β cyclodextrin although large differences are found between α - and β -anomers, the former exhibiting the larger solubility. The inclusion capacity of the these derivs. has been investigated using NMR spectroscopy as the major anal. technique for various host-quest pairs. The apparent

discrepancies between the intrinsic solubilities of these host mols. and their ability to solubilize hydrophobic hosts can be explained from geometrical considerations derived from detailed NMR studies. The resp. roles of the side of inclusion, of steric effects and of stabilizing

interactions are evidenced and allow an a priori selection of the optimal host derivative for a given guest mol.

L11 ANSWER 32 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1995:421701 CAPLUS <<LOGINID::20080328>>

DOCUMENT NUMBER:

Incorporation of sulfonated cyclodextri into polypyrrole: an approach for the electro-controlled delivering of neutral drugs

AUTHOR(S): Bidan, G.; Lopez, C.; Mendes-Viegas, F.; Vieil, E.;

Gadelle, A. Lab. Electrochimie Moleculaire, Centre Etudes CORPORATE SOURCE:

Nucleaires Grenoble, Grenoble, 38054, Fr. Biosensors & Bioelectronics (1995), 10(1/2), 219-29

CODEN: BBIOE4; ISSN: 0956-5663 PUBLISHER: Elsevier Advanced Technology

DOCUMENT TYPE:

LANGUAGE: English

The electro-controlled delivery of drugs based on the doping-dedoping mechanism of Electro-Conducting Polymers is restricted to charged substances acting as dopants. In order to overcome this limitation, this study presents an approach where the trapping/delivering is based on host-quest interaction. As an example of a neutral quest, the mol. N-methylphenothiazine (NMP) is encapsulated in the host, heptasulfonated β - cyclodextrin (β -CDSO3-), which is tailor-made to dope polypyrrole (PPy). The original synthetic method for β -CDS03is based on sulfonation of the periodated β -CD in the phase transfer medium. As a consequence of their size and of their multicharged character, β -CDS03- \hat{s} are fixed dopants. The stability of the β -CDSO3- entrapment is checked by Optical Beam Deflection (mirage effect) measurements. The ionic movements associated with the switching of the β-CDS03- doped PPy (PPy+, β-CDS03-) film appear to be mainly due to cations with this technique. Cyclic voltammetry expts. confirm the

entrapment of neutral NMP by simply dipping the PPY+, β -CDSO3- film in a CH3CN solution containing NMP. Repeated electrochem. cycling of such a reservoir electrode indicates the progressive elimination of NMP from the (PPy+, β-CDSO3- [NMP]) film.

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ACCESSION NUMBER:
                           1995:316135 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                           Conductive polymer doped with sulfonated
                           cyclodextrin salt and device for capturing and/or delivering an active substance using this
                           polymer.
                           Vieil, Eric; Bidan, Gerard; Gadelle, Andree;
                           Mendes, Viegas Maria-Fatima
PATENT ASSIGNED(S):
                           Commissariat a l'Energie Atomique, Fr.
                           Eur. Pat. Appl., 10 pp.
                           CODEN: EPXXDW
DOCUMENT TYPE:
                           Patent
LANGUAGE:
                           French
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                           KIND
                                  DATE
                                               APPLICATION NO.
                                                                        DATE
     EP 627747
                           A1
                                   19941207
                                               EP 1994-401204
                                                                        19940601
         R: CH, DE, FR, GB, IT, LI
     FR 2706067
                           A1
                                               FR 1993-6655
                                                                        19930603
     FR 2706067
     US 5480924
                                   19960102
                                               US 1994-246125
                                                                        19940519
                            A
                                               JP 1994-122727
                                                                        19940603
     US 5587466
                           A
                                  19961224
                                               US 1995-539437
                                                                        19951005
PRIORITY APPLN. INFO.:
                                               FR 1993-6655
                                                                     A 19930603
                                               US 1994-246125
                                                                     A3 19940519
                          MARPAT 122:94365
    In a conductive polymer doped by a sulfonated cyclodextrin salt
and a device for capturing and/or delivering an active substance using
     this polymer, the dopant has formula I, in which n is 2-50, M+ is Na+,
     Li+, K+, Mg+1/2 or NH4+ and R is -SO3M+ or -OH, R being different from the
     ring of the other. The doped conductive polymer can be used as the active
     electrode in an electrochem. device.
L11 ANSWER 34 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                           1995:253021 CAPLUS <<LOGINID::20080328>>
DOCUMENT NUMBER:
                           122:187960
                           Synthesis of cyclohexakis- and cycloheptakis-
                           (1\rightarrow 4)-(7-amino-6,7-dideoxy-\alpha-D-gluco-
                           heptopyranosyl), homoanalogues of 6-amino-6-deoxy-
                           cyclomaltooligosaccharides
AUTHOR(S):
                           Defaye, Jacques; Gadelle, Andree
CNRS and CEA, Departement de Recherche Fondamentale
CORPORATE SOURCE:
                           sur la Matiere Condensee/SESAM, Centre d'Etudes de
                           Grenoble, Grenoble, F-38054, Fr.
                           Carbohydrate Research (1994), 265(1), 129-32
                           CODEN: CRBRAT; ISSN: 0008-6215
PUBLISHER:
                           Elsevier
DOCUMENT TYPE:
                           Journal
LANGUAGE:
                           English
OTHER SOURCE(S):
                           CASREACT 122:187960
   Aminodideoxycyclodextrins I (n = 6, 7) were prepared from
     iododeoxycylodextrins via cyanation and reduction
L11 ANSWER 35 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN
                           1994:157132 CAPLUS <<LOGINID::20080328>>
ACCESSION NUMBER:
DOCUMENT NUMBER:
                           120:157132
                           Nuclear magnetic resonance study of a polar headgroup
                           determined \alpha- \underline{\text{cyclodextrin}}\text{-phospholipid}
                           association
AUTHOR(S):
                           Fauvelle, F.; Debouzy, J. C.; Nardin, R.;
                           Gadelle, A. Unite de Biophysique, CRSSA, La Tronche-Grenoble, Fr.
CORPORATE SOURCE:
                           Bioelectrochemistry and Bioenergetics (1994), 33(1),
                           95-9
                           CODEN: BEBEBP; ISSN: 0302-4598
DOCUMENT TYPE:
LANGUAGE:
                           English
AB In order to investigate the hemolytic activity of α-
     cyclodextrin, the interactions of this cyclic oligosaccharide with
```

selected membrane phospholipids were studied by 1H-NMR and 31P-NMR. Two

natural phospholipids differing by their polar headgroup, phosphatidylcholine and phosphatidylinositol, were tested. The results suggest that interactions of α - cyclodextrin with phospholipids are at least modulated by the nature of the polar headgroup in a first step. The acyl chains could be implicated in a second step.

L11 ANSWER 36 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1991:82321 CAPLUS <<LOGINID::20080328>>

DOCUMENT NUMBER:

Selective halogenation of cyclic maltose

oligosaccharides in the C-6 position and synthesis of per (3,6-anhydro) cyclic maltose oligosaccharides

AUTHOR(S):

Gadelle, Andree; Defaye, Jaques Dep. Rech. Fondam., Cent. Etud. Nucl. Grenoble, CORPORATE SOURCE:

Grenoble, F-38041, Fr.

Angewandte Chemie (1991), 103(1), 94-5 (See also Angew. Chem., Int. Ed. Engl., 1991, 30(1), 78-80)

CODEN: ANCEAD; ISSN: 0044-8249

DOCUMENT TYPE:

German LANGUAGE:

AB Cyclic maltose oligosaccharides were treated with PPh3 and iodine (or

bromine) to give the per-6-deoxy-6-halo derivs. Treatment of per(6-deoxy-6-iodo) cyclic maltose oligosaccharide with aqueous NaOH gave the per(3,6-anhydro) derivs.

L11 ANSWER 37 OF 37 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1990:179645 CAPLUS <<LOGINID::20080328>>

112:179645 DOCUMENT NUMBER:

Stereoselective thioglycoside synthesis. Part X.

Branched thiocyclomalto-oligosaccharides: synthesis

and properties of $6-S-\alpha-$ and

6-S-β-D-glucopyranosyl-6-thiocyclomaltoheptaose

Defaye, Jacques; Gadelle, Andree; Guiller, Alain; Darcy, Raphael; O'Sullivan, Thomas AUTHOR(S):

CORPORATE SOURCE: Dep. Rech. Fondam., Cent. Etud. Nucl., Grenoble,

Carbohydrate Research (1989), 192, 251-8

CODEN: CRBRAT; ISSN: 0008-6215

DOCUMENT TYPE: Journal LANGUAGE: English

CASREACT 112:179645

 $6-S-\alpha-$ (I) And $6-S-\beta-D-$ glucopyranoysl-6-thiocyclomaltoheptaose (II) have been prepared by treatment of 6-0-p-tolylsulfonylcyclomaltoheptaos

e with the sodium salts of 1-thio- α - and $-\beta$ -D-glucopyranose, resp., in 1,3-dimethyl-2-oxohexahydropyrimidine. Compds. I and II are

more soluble in water than cyclomaltoheptaose and enhance the solubility of hydrophobic compds. by inclusion.